RAW SEQUENCE LISTING

The Biotechnology Systems Branch of the Scientific and Technical Information Center (STIC) no errors detected.

| Application Serial Number: | 10/536, 677A |
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| Source: | TFW16 |
| Date Processed by STIC: | 3/23/2007 |

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IFW16

RAW SEQUENCE LISTING DATE: 03/23/2007
PATENT APPLICATION: US/10/536,677A TIME: 14:46:51

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4 <110> APPLICANT: Clark, Georgina Jane
             Hart, Derek Nigel Jo
     7 <120> TITLE OF INVENTION: THERAPEUTIC AND DIAGNOSTIC AGENTS
     10 <130> FILE REFERENCE: DAVI257.001APC
     12 <140> CURRENT APPLICATION NUMBER: US 10/536,677A
C--> 13 <141> CURRENT FILING DATE: 2006-06-05
     15 <150> PRIOR APPLICATION NUMBER: PCT/AU2003/001586
     16 <151> PRIOR FILING DATE: 2003-11-28
     18 <150> PRIOR APPLICATION NUMBER: AU 2002952993
    19 <151> PRIOR FILING DATE: 2002-11-29
    22 <160> NUMBER OF SEQ ID NOS: 28
    25 <170> SOFTWARE: PatentIn version 3.1
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   · 32 <212> TYPE: DNA
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    42 acaaggaagc agagagcaga agaaaagcag aagcgaagct cagatctgct gggaggaaga
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    44 ttacattttg tcccctcctg gggtcttgca cagtggcagg tgacattcgt gttacaggaa
    46 tgactgccag ggcctgggcc tcgtggcggt cttcagctct gctcctcctg cttgtcccag
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    48 gctattttcc tctgagccac cccatgaccg tggcgggccc cgtgggggga tccctgagtg
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    50 tgcagtgteg ctatgagaag gaacacagga ccctcaacaa attctggtgc agaccaccac
                                                                             420
    52 agatteteeg atgtgacaag attgtggaga ceaaagggte ageagggaaa aggaatggee
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    54 gagtgtccat cagggacagt cctgcaaacc tcagcttcac agtgaccctg gagaatctca
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    56 cagaggagga cgcaggcacc tactggtgtg gggtggatac accgtggctc cgagactttc
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    58 atgateceat tgtegaggtt gaggtgteeg tgtteeegge egggaegaee acageeteea
                                                                             660
    60 gececeagag etecatggge aceteaggte eteceaegaa getgeeegtg cacacetgge
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    62 ccagegtgae cagaaaggae ageeeegaae eeageeeaca eeetggetee etgtteagea
                                                                             780
    64 atgtccgctt cctgctcctg gtcctcttgg agctgccct gctcctgagc atgctgggtg
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    66 ccgtcctctg ggtgaacaga cctcagagaa gctctagaag caggcagaat tggcccaagg
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    68 gtgagaacca gtagcatctg ctgtccatca aggccctgtg ctgcaacaga gcccctctgg
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    70 ggactggaat gacctcctga ccatcaaggc ctgcaacaga gcccctctgg gggactggaa
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    72 tgacctcctg accactccct cccgggctgc tctctccaac atctcctgga atcctttgtg
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    85 <213 > ORGANISM: Homo sapiens
    88 <400> SEQUENCE: 2
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TIME: 14:46:51

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106 Cys Asp Lys Ile Val Glu Thr Lys Gly Ser Ala Gly Lys Arg Asn Gly
107 65
110 Arg Val Ser Ile Arg Asp Ser Pro Ala Asn Leu Ser Phe Thr Val Thr
114 Leu Glu Asn Leu Thr Glu Glu Asp Ala Gly Thr Tyr Trp Cys Gly Val
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                                     105
118 Asp Thr Pro Trp Leu Arg Asp Phe His Asp Pro Ile Val Glu Val Glu
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                                 120
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122 Val Ser Val Phe Pro Ala Gly Thr Thr Ala Ser Ser Pro Gln Ser
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                            135
126 Ser Met Gly Thr Ser Gly Pro Pro Thr Lys Leu Pro Val His Thr Trp
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130 Pro Ser Val Thr Arg Lys Asp Ser Pro Glu Pro Ser Pro His Pro Gly
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134 Ser Leu Phe Ser Asn Val Arg Phe Leu Leu Leu Val Leu Leu Glu Leu
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138 Pro Leu Leu Ser Met Leu Gly Ala Val Leu Trp Val Asn Arg Pro
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160 ggaccetcaa caaatactgg tgcagaccac cacagatttt cetatgtgac aagattgtgg
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162 agaccaaagg gtcagcagga aaaaggaacg gccgagtgtc catcagggac agtcctgcaa
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164 acctcagett cacagtgace etggagaate teacagagga ggatgeagge acctaetggt
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166 gtggggtgga tacaccgtgg ctccgagact ttcatgatcc cgttgtcgag gttgaggtgt
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168 ccgtgttccc ggcatcaacg tcaatgacac ctgcaagtat cactgcggcc aagacctcaa
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170 caatcacaac tgcatttcca cctgtatcat ccactaccct gtttgcagtg ggtgccaccc
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172 acagtgccag catccaggag gaaactgagg aggtggtgaa ctcacagctc ccgctgctcc
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174 tetecetget ggeattgttg etgettetgt tggtggggge etecetgeta geetggagga
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176 tgtttcagaa atggatcaaa gctggtgacc attcagagct gtcccagaac cccaaqcagq
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178 ctgccacgca gagtgagctg cactacgcaa atctggagct gctgatgtgg cctctgcagg
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180 aaaagccagc accaccaagg gaggtggagg tggaatacag cactgtggcc tcccccaggg
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182 aagaacttca ctatgcctcg gtggtgtttg attctaacac caacaggata gctgctcaga
                                                                          840
184 ggcctcggga ggaggaacca gattcagatt acagtgtgat aaggaagaca taggcttttg
                                                                          900
186 tectgeeteg ceateggage teteatggge eccaggaagt ceagggacag etceettata
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188 cctggcccac gtccttctca gcctgccctc gacaacagtg accaacagac aggcagctgg
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                                                                          1140
194 cccctttgcc cctgcttcat cccagctctg tgtgtggagg acaaagcttc ttcctgcgtg
                                                                          1200
196 gctccaggaa aagatgtggc tcacgtaggt ggcacctgcc aatagctttg tcaatcacag
198 ccccatagga acgtctggaa ttgcttggga gttggggaga actgtcaaga agagtgaaga
200 gagtgccaaa gcggagatct gttcacctgg gggccatgga gggqqgaccc actaaaqatc
                                                                         1380
202 aagatcaaag atteteecca teteacagae aaggaaactq aqqecaqaqq qaqqaqaa
                                                                          1440
20.4 ttgctcatgg ctccagaact ggtggcaagt ttctctggac tcttaggttt atttttaata
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224 Ala Leu Ser Lys Cys Arg Thr Val Ala Gly Pro Trp Gly Ser Leu Ser
                20
                                     25
228 Val Sln Cys Pro Tyr Slu Lys Slu His Arg Thr Leu Asn Lys Tyr Trp
            35
232 Cys Arg Pro Pro Gln Ile Phe Leu Cys Asp Lys Ile Val Glu Thr Lys
236 Gly Ser Ala Gly Lys Arg Asn Gly Arg Val Ser Ile Arg Asp Ser Pro
237 65
                        70
                                             75
240 Ala Asn Leu Ser Phe Thr Val Thr Leu Glu Asn Leu Thr Glu Glu Asp
244 Ala Gly Thr Tyr Trp Cys Gly Val Asp Thr Pro Trp Leu Arg Asp Phe
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                                    105
248 His Asp Pro Val Val Glu Val Glu Val Ser Val Phe Pro Ala Ser Thr
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                                120
252 Ser Met Thr Pro Ala Ser Ile Thr Ala Ala Lys Thr Ser Thr Ile Thr
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                            135
                                                 140
256 Thr Ala Phe Pro Pro Val Ser Ser Thr Thr Leu Phe Ala Val Gly Ala
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260 Thr His Ser Ala Ser Ile Gln Glu Glu Thr Glu Glu Val Val Asn Ser
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264 Gln Leu Pro Leu Leu Leu Ser Leu Leu Ala Leu Leu Leu Leu Leu Leu
                                    185
268 Val Gly Ala Ser Leu Leu Ala Trp Arg Met Phe Gln Lys Trp Ile Lys
269
            195
                                200
272 Trp Ile Lys Ala Gly Asp His Ser Glu Leu Ser Gln Asn Pro Lys Gln
                            215
                                                 220
276 Ala Ala Thr Gln Ser Glu Leu His Tyr Ala Asn Leu Glu Leu Leu Met
                        230
                                            235
280 Trp Pro Leu Gln Glu Lys Pro Ala Pro Pro Arg Glu Val Glu Val Glu
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284 Tyr Ser Thr Val Ala Ser Pro Arg Glu Glu Leu His Tyr Ala Ser Val
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288 Val Phe Asp Ser Asn Thr Asn Arg Ile Ala Ala Gln Arg Pro Arg Glu
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PATENT APPLICATION: US/10/536,677A TIME: 14:46:51

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310 gctctgtgac tggcactgcg ggggactctc tgacagtgtg gtgtcagtat gagagcatgt
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312 acaagggata taacaagtac tggtgccgag gacagtacga cacgtcatgt gagagcattg
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314 tggagaccaa gggagaagag aaggtggaga ggaatggccg cgtgtccatc agagaccacc
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316 cggaggetet egeetteact gtgaccatge agaaceteaa tgaagatgat getggatett
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318 actggtgcaa aattcagaca gtgtgggtcc tggattcatg gtcacgcgat ccctcggacc
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320 tggttagggt gtatgtttcc ccagcaatta caaccccaag gaggaccaca catccagcca
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324 cccaaaattc agggttccgg ctcagcagcc ctcacttcct gctcgtggtc cttctgaagc
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338 <213> ORGANISM: Homo sapiens
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355 Trp Cys Arg Gly Gln Tyr Asp Thr Ser Cys Glu Ser Ile Val Glu Thr
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359 Lys Gly Glu Glu Lys Val Glu Arg Asn Gly Arg Val Ser Ile Arg Asp
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363 His Pro Glu Ala Leu Ala Phe Thr Val Thr Met Gln Asn Leu Asn Glu
364
367 Asp Asp Ala Gly Ser Tyr Trp Cys Lys Ile Gln Thr Val Trp Val Leu
368
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                                     105
371 Asp Ser Trp Ser Arg Asp Pro Ser Asp Leu Val Arg Val Tyr Val Ser
                                120
375 Pro Ala Ile Thr Thr Pro Arg Arg Thr Thr His Pro Ala Thr Pro Pro
                            135
379 Ile Phe Leu Val Val Asn Pro Gly Arg Asn Leu Ser Thr Arg Glu Val
                        150
                                             155
383 Leu Thr Gln Asn Ser Gly Phe Arg Leu Ser Ser Pro His Phe Leu Leu
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                                         170
387 Val Val Leu Leu Lys Leu Pro Leu Leu Leu Ser Met Leu Gly Ala Val
388
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415 gatgctgaca gttattggtg tgggactgag agacctggaa ttgatcttgg ggtcaaagtt
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417 caagtgacca ttaacccagc tcagtgcctg agtctgttgc ccacagatga cagggtgatg
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419 gttccagttt cagcccacag gccaaaggga cccccttccc tggtaaccag agaccccaat
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451 Cys Asn Ile Leu Val Lys Thr Asn Gly Ser Glu Gln Glu Val Lys Lys
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                                             75
455 Asn Arg Val Ser Ile Arg Asp Asn Gln Lys Asn His Val Phe Thr Val
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459 Thr Met Glu Asn Leu Lys Arg Asp Asp Ala Asp Ser Tyr Trp Cys Gly
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463 Thr Glu Arg Pro Gly Ile Asp Leu Gly Val Lys Val Gln Val Thr Ile
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467 Asn Pro Ala Gln Cys Leu Ser Leu Leu Pro Thr Asp Asp Arg Val Met
468
                            135
                                                 140
471 Val Pro Val Ser Ala His Arg Pro Lys Gly Pro Pro Ser Leu Val Thr
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VERIFICATION SUMMARY

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